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DESCRIPTION

HIGH STRENGTH STAINLESS STEEL PIPE FOR USE IN OIL WELL HAVING SUPERIOR CORROSION RESISTANCE AND MANUFACTURING METHOD THEREOF

Technical Field

gas wells. In particular, the present-invention relates to a high strength stainless steel having superior corrosion resistance, which is suitably used in an oil well and gas well in a very severe corrosion environment containing carbon dioxide (CO₂), chloride ions (Cl⁻), and the like. In the present-invention, the The "high strength stainless steel pipe" indicates a stainless steel pipe having a yield strength of 654 MPa (95 ksi) or more.

Background-Art

[0002] In recent years, in response to steep rise in crude oil price and to depletion of petroleum oil resources anticipated in the near future, deeper oil fields, which have not be taken into consideration in the past, very corrosive sour gas fields, the development of which was abandoned once in the past, and the like have been aggressively developed on a worldwide basis. The oil fields and gas fields as described above are generally located in very deep places, and in addition, these oil and gas fields are in a very severe corrosive environment in which the temperature is high and CO₂, Cl⁻, and the like are present. Hence, as an oil-well steel pipe used for mining oil and gas fields as described above, a steel pipe having high strength and also having superior corrosion resistance is required.

[0003] Heretofore, in oil wells and gas wells in an environment containing CO₂, Cl⁻, and the like, 13%Cr martensite stainless steel pipes, which have superior CO₂ corrosion resistance, have been generally used as an oil-well steel pipe. However, there has been a problem in that a general martensite stainless steel cannot withstand the use in an environment in which a large amount of Cl⁻ is present and the temperature is high, such as more than 100°C. Hence, in a well in which steel pipes and the like are required to have corrosion resistance, a dual phase stainless steel pipe has been used. However, since the dual phase stainless steel pipe contains a large amount of alloy elements, hot workability thereof is not superior, and hence a specific hot working can only be used for forming the dual phase stainless steel pipe, thereby causing the increase in cost. In addition, when the yield strength of a conventional 13%Cr martensite stainless steel pipe is more than 654 MPa, the toughness thereof is seriously degraded, and hence there has been a problem in that the 13%Cr martensite stainless steel pipe may not be used.

[0004] In addition, in recent years, development of oil wells in a-cold regions has been increasingly carried out, and hence. Hence, besides high strength, superior low-temperature toughness has also been required for the steel pipe in many cases.

[0005] According to the situations as-described above, a high strength 13%Cr martensite stainless steel pipe for use in oil wells has been strongly desired, which is primarily formed of inexpensive 13%Cr martensite stainless steel having excellent hot workability and which has a high yield strength of more than 654 MPa (95 ksi), superior CO₂ corrosion resistance, and a high toughness.

[0006] In response to the requirements described above, for example, in Patent Documents 1, 2, 3, 4, and 5 Japanese Unexamined Applications 8-120345, 9-268349 and 10-1755 and Japanese Patents 28-14528 and 32-51648, improved martensite stainless steel or a steel pipe thereof have

been proposed which are obtained by improving the corrosion resistance of 13%Cr martensite stainless steel or a steel pipe thereof.

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A technique disclosed in Patent Document 1 Japanese Unexamined Application 8-[0007] 120345 is a method for manufacturing a martensite stainless steel seamless pipe having superior corrosion resistance. According to the method described above, after a 13%Cr stainless-steel raw material having a composition in which the content of C is controlled in the range of 0.005% to 0.05%, 2.4% to 6% of Ni and 0.2% to 4% of Cu are collectively added, 0.5% to 3% of Mo is further added, and a Nieq is adjusted to 10.5 or more is processed by hot working, cooling at a rate faster than that of air cooling is performed. In addition, alternatively, heating Heating may further be performed to a temperature in the range of (the Ac₃ transformation point + 10°C) to (the Ac₃ transformation point + 200°C) or may further be performed to a temperature in the range of the Ac₁ transformation point to the Ac₃ transformation point, followed by cooling to room temperature at a cooling rate faster than that of air cooling, so that tempering is performed. According to the technique described in Patent Document 1, it is said that Japanese Unexamined Application 8-120345, a martensite stainless steel seamless pipe can be manufactured which simultaneously has a high strength equivalent to or more than that of API-C95 grade, corrosion resistance in an environment at 180°C or more containing CO₂, and the SCC resistance.

[0008] A technique disclosed in Patent Document 2Japanese Unexamined Application 9-268349 is a method for manufacturing a martensite stainless steel having superior resistance to sulfide stress cracking. According to the method described above, after 13%Cr martensite stainless steel having a composition in which 0.005% to 0.05% of C and 0.005% to 0.1% of N are contained, and in which Ni, Cu, and Mo are controlled in the ranges of 3.0% to 6.0%, 0.5% to 3% and 0.5% to 3%, respectively, is processed by hot working, followed by spontaneous

cooling to room temperature, heating is performed to a temperature in the range of (the Ac₁ point + 10°C) to (the Ac₁ point + 40°C), and the stainless steel is held for 30 to 60 minutes at that temperature and is then cooled to a temperature to the Ms point or less. Subsequently, tempering is performed at a temperature of the Ac₁ point or less, so that a texture is formed in which tempered martensite and 20 percent by volume or more of a γ phase are both present. According to the technique described in Patent Document 2, it is said that since aA tempered martensite texture containing 20 percent by volume or more of a γ phase is formed, the resistance to sulfide stress cracking is significantly improved.

Application 10-1755, martensite stainless steel has a composition containing 10% to 15% of Cr in which the content of C is controlled in the range of 0.005% to 0.05%, 4.0% or more of Ni and 0.5% to 3% of Cu are collectively added, 1.0% to 3.0% of Mo is further added, and in addition, the Nieq is controlled to -10 or more. By performing tempering, a texture is formed containing a tempered martensite phase, a martensite phase, and a retained austenite phase so that the total fraction of the tempered martensite phase and the martensite phase is set to 60% to 90%, thereby obtaining martensite stainless steel having superior corrosion resistance and resistance to sulfide stress cracking. According to the technique described in Patent Document 3, it is said that the The corrosion resistance and the resistance to sulfide stress cracking in a wet carbon dioxide gas environment and in a wet hydrogen sulfide environment are improved.

[0010] A technique described in <u>Japanese</u> Patent <u>Document 428-14528</u> relates to a martensite stainless steel material for use in oil wells, having superior resistance to sulfide stress cracking, the stainless steel material having a steel composition in which more than 15% to 19% or Cr is contained, 0.05% or less of C, 0.1% or less of N, and 3.5% to 8.0% of Ni are contained,

and 0.1% to 4.0% of Mo is further contained, and in which 30Cr+36Mo+14Si-28Ni≤455 (%) and 21Cr+25Mo+17Si+35Ni≤731 (%) are simultaneously satisfied. According to the technique described in Patent Document 4, it is said that aA steel material having superior corrosion resistance in a severe oil well environment in which chloride ions, a carbon dioxide gas, and a small amount of a hydrogen sulfide gas are present.

[0011] A technique described in <u>Japanese</u> Patent <u>Document 532-51648</u> relates to a precipitation hardened martensite stainless steel having superior strength and toughness, the stainless steel having a steel composition in which 10.0% to 17% or Cr is contained, 0.08% or less of C, 0.015% or less of N, 6.0% to 10.0% of Ni, and 0.5% to 2.0% of Cu are contained, and 0.5% to 3.0% of Mo is further contained, and having a texture in which, owing to a cold working of 35% or more and annealing, the average crystal particle diameter is set to 25 μm or less and the number of precipitates, which are precipitated in a matrix and which have a particle diameter of 5×10⁻² μm or more, is reduced to 6×10⁶/mm² or less. According to the technique described in Patent Document 5, it is said that since Since a texture is formed containing fine crystal particles and having a small amount of precipitates, precipitation hardened martensite stainless steel, which has a high strength and causes no decrease in toughness, can be provided.

Patent Document 1: Japanese Unexamined Patent Application Publication No. 8-120345

Patent Document 2: Japanese Unexamined Patent Application Publication No. 9-268349

Patent Document 3: Japanese Unexamined Patent Application Publication No. 10-1755

Patent Document 4: Japanese Patent No. 2814528

Patent Document 5: Japanese Patent No. 3251648

Disclosure of Invention

However, there has been a problem in that improved 13%Cr martensite stainless steel pipes manufactured by the techniques disclosed in Patent Documents 1, 2, 3, 4, and 5 discussed above cannot stably exhibit desired corrosion resistance in a severe corrosive environment in which CO₂, Cl⁻, and the like are present and the temperature is high, such as more than 180°C.

Summary

[0013] The present-invention was made in consideration of the conventional techniques described above. An object Aspects of the present this invention is to provide a high strength stainless steel pipe for use in oil wells and the manufacturing method thereof, the high strength stainless steel pipe being inexpensive, and having superior hot workability, a high yield strength of more than 654 MPa, and superior corrosion resistance such as superior CO₂ corrosion resistance even in a severe corrosive environment in which CO₂, Cl⁻ and the like are present and the temperature is high, such as up to 230°C.

[0014] In order to achieve the object described above, intensive research on various factors relating to the hot workability and corrosion resistance was carried out by the inventors of the present invention.

[0015] — In manufacturing a conventional martensite stainless steel seamless pipe, when a martensite single-phase is not obtained due to the formation of a ferrite phase, the strength is decreased and the hot workability is degraded; hence it has been generally believed that manufacturing of the steel pipe cannot be easily performed. Accordingly, as disclosed in Japanese Unexamined Patent Application Publication No. 8 246107, generally in a 13%Cr stainless steel seamless pipe for use in oil wells, for manufacturing, the composition thereof has

been controlled so that the formation of ferrite is suppressed to obtain a texture formed of a martensite single phase.

[0016] Accordingly, intensive research on the influences of components on the hot workability was further carried out in detail by the inventors of the present invention. As a result, it was found that when the steel composition is controlled to satisfy the following equation (2), the hot workability is significantly improved, and that generation of crack in hot working can be prevented.

(where Cr, Ni, Mo, Cu, C, Si, Mn, and N represent respective contents on a mass percent basis)

[0017] Fig. 1 shows the relationship between the value of the left hand side of the equation

(2) and the length of crack-generated in an end surface of a 13%Cr stainless steel seamless pipe
in hot working (that is, in pipe making of a seamless steel pipe). As can be seen from Fig. 1, it is
understood that when the value of the left hand side of the equation (2) is 8.0 or less, or the left
hand side of the equation (2) is 11.5 or more and is preferably 12.0 or more, the generation of
crack can be prevented. A value of the left hand side of the equation (2) of 8.0 or less represents
a region in which ferrite is not formed at all, and this region corresponds to a region defined by
the conventional concept of improvement in hot workability in which the formation of a ferrite
phase is not allowed. In addition, as the value of the left hand side of the equation (2) is
increased, the amount of ferrite thus formed is increased, and in the region in which the value of
the left hand side is 11.5 or more, a relatively large amount of ferrite is formed. That is, the
inventors of the present invention first found that when the concept is employed that is totally
different from the conventional one in the past, that is, when the composition is adjusted to have

a value of the left hand side of 11.5 or more so that a texture containing a relatively large amount of ferrite is used in pipe making, the hot workability can be significantly improved.

[0018] Fig. 2 shows the relationship between the amount of ferrite and the length of crack generated in the end surface of a 13%Cr stainless steel seamless pipe in hot working, the relationship being obtained based on the data described above. As can be seen from Fig. 2, as is the conventional concept, cracks are not generated when the amount of ferrite is 0 percent by volume; however, as ferrite is formed, cracking starts to occur. However, when the amount of ferrite is further increased to 10 percent by volume or more and preferably 15 percent by volume or more, the generation of cracks can be prevented, and this phenomenon is totally different from that based on the conventional concept. That is, when the components are adjusted to satisfy the equation (2), and a ferrite-martensite dual phase is formed in which an appropriate amount of a ferrite phase is formed, the hot workability is improved, and the generation of cracks can be prevented. In addition, it was also found that when a ferrite martensite dual phase texture is used, a strength required for oil well pipes can also be ensured.

[0019] However, when the components are adjusted to satisfy the equation (2) so as to form a ferrite-martensite dual phase texture, the corrosion resistance may be degraded in some cases due to the distribution of elements which occurs during heat treatment. When the dual phase texture is formed, since elements such as C, Ni, and Cu forming an austenite phase are diffused to a martensite phase, and elements such as Cr and Mo forming a ferrite phase are diffused to a ferrite phase, as a result, variation in component between the phases occurs in a final product obtained after heat treatment. In the martensite phase, since the amount of Cr effective for corrosion resistance is decreased, and the amount of C degrading corrosion resistance is

increased, as a result, the corrosion resistance may be degraded in some cases as compared to that of a uniform texture.

[0020] Accordingly, intensive research on the influences of components on the corrosion resistance was carried out by the inventors of the present invention. Consequently, it was found that by adjusting components to satisfy the following equation (1), even when a ferrite austenite dual phase texture is formed, sufficient corrosion resistance can be ensured.

$$Cr + 0.65Ni + 0.6Mo + 0.55Cu - 20C \ge 19.5$$
 (1)

(where Cr, Ni, Mo, Cu, and C represent the respective contents on a mass percent basis.)

[0021] Fig. 3 shows the relationship between the value of the left hand side of the equation (1) and the corrosion rate in a high temperature environment at 230°C containing CO₂ and Cl⁻. As can be seen from Fig. 3, by adjusting the components to satisfy the equation (1), even when a ferrite austenite dual phase texture is formed, in a high temperature environment at 230°C containing CO₂ and Cl⁻, sufficient corrosion resistance can be ensured.

[0022] — As apparent from the equation (1), in order to improve the corrosion resistance, the content of Cr is advantageously increased. However, Cr promotes the formation of ferrite. Hence, in order to suppress the formation of ferrite, Ni in an amount corresponding to the content of Cr was necessary to be added in the past. However, when the content of Ni is increased so as to correspond to the content of Cr, an austenite phase is stabilized, and as a result, a problem may arise in that a strength required for oil well pipes cannot be ensured.

[0023] In order to solve this problem, the inventors of the present invention found that when the content of Cr is increased while a ferrite austenite dual phase texture containing an appropriate amount of a ferrite phase is maintained, a remaining amount of an austenite phase can be reduced and a sufficient strength as an oil well pipe can be ensured.

[0024] Fig. 4 shows the relationship between the content of Cr and the yield strength YS of a 13%Cr stainless steel seamless pipe containing a ferrite austenite dual phase texture processed by heat treatment, the relationship being obtained by the inventors of the present invention. In Fig. 4, the relationship between the content of Cr and the yield strength YS of a martensite single phase texture or a martensite austenite dual phase texture processed by heat treatment is also shown. From Fig. 3, it was first found that when the ferrite austenite dual phase texture containing an appropriate amount of a ferrite phase is maintained, and the content of Cr is increased, a sufficient strength as an oil well pipe can be ensured. On the other hand, when the texture is a martensite single phase or a martensite-austenite dual phase texture, as the amount of Cr is increased, the YS is decreased.

[0025] Research was further carried out based on the above findings, and as a result, the present invention was finally made. That is, the present invention includes the following.

There is provided a high strength stainless steel pipe for use in oil wells, which has superior corrosion resistance, comprising on a mass percent basis: about 0.005% to about 0.05% of C; about 0.05% to about 0.5% of Si; about 0.2% to about 1.8% of Mn; about 0.03% or less of P; about 0.005% or less of S; about 15.5% to about 18% of Cr; about 1.5% to about 5% of Ni; about 1% to about 3.5% of Mo; about 0.02% to about 0.2% of V; about 0.01% to about 0.15% of N; about 0.006% or less of O; and the balance being Fe and unavoidable impurities, in which the following equations (1) and (2) are satisfied:

$$Cr+0.65Ni+0.6Mo+0.55Cu-20C \ge 19.5$$
 (1)

$$Cr+Mo+0.3Si-43.5C-0.4Mn-Ni-0.3Cu-9N\geq11.5$$
 (2)

(where Cr, Ni, Mo, Cu, C, Si, Mn, and N represent the respective contents on a mass percent basis).

- (2) According to the above (1), in In addition to the above composition, the high strength stainless steel pipe for use in oil wells may further comprises about 0.002% to about 0.05% of Al on a mass percent basis.
- (3) According to the above (1) or (2), in the high strength stainless steel pipe for use in oil-wells, the The content of C is may be in the range of about 0.03% to about 0.05% on a mass percent basis.
- (4) According to one of the above (1) to (3), in the high strength stainless steel-pipe for use in oil wells, the The content of Cr is may be in the range of about 16.6% to less than about 18% on a mass percent basis.
- (5) According to one of the above (1) to (4), in the high strength stainless steel pipe for use in oil wells, the The content of Mo is may be in the range of about 2% to about 3.5% on a mass percent basis.
- (6) According to one of the above (1) to (5), in addition to the above composition, the The high strength stainless steel pipe for use in oil wellsmay further comprises about 3.5% or less of Cu on a mass percent basis.
- (7) According to the above (6), in the high strength stainless steel pipe for use in oil wells, the The content of Cu is may be in the range of about 0.5% to about 1.14% on a mass percent basis.
- (8) According to one of the above (1) to (7), in addition to the above composition, the The high strength stainless steel pipe for use in oil wellsmay further comprises at least one element selected from the group consisting of about 0.2% or less of Nb, about 0.3%

or less of Ti, <u>about 0.2%</u> or less of Zr, <u>about 3%</u> or less of W, and <u>about 0.01%</u> or less of B on a mass percent basis.

- (9) According to one of the above (1) to (8), in In addition to the above composition, the high strength stainless steel pipe for use in oil wells may further comprises about 0.01% or less of Ca on a mass percent basis.
- (10) According to one of the above (1) to (9), the The high strength stainless steel pipe for use in oil wells has may have a texture containing a martensite phase as a primary phase and a ferrite phase at a volume fraction of about 10% to about 60%.
- (11) According to the above (10), in the high-strength-stainless steel pipe for use in-oil wells, the The ferrite phase has may have a volume fraction of about 15% to about 50%.
- (12) According to the above (10) or (11), in the high strength stainless steel pipe for use in oil wells, the The texture may further contains an austenite phase at a volume fraction of about 30% or less.
- (13) There is provided a method for manufacturing a high strength stainless steel pipe for use in oil wells having superior corrosion resistance, comprising the steps of: preparing a steel pipe raw material which contains on a mass percent basis, about 0.005% to about 0.05% of C; about 0.05% to about 0.5% of Si; about 0.2% to about 1.8% of Mn; about 0.03% or less of P; about 0.005% or less of S; about 15.5% to about 18% of Cr; about 1.5% to about 5% of Ni; about 1% to about 3.5% of Mo; about 0.02% to about 0.2% of V; about 0.01% to about 0.15% of N; about 0.006% or less of O; and the balance being Fe and unavoidable impurities, and which satisfies the following equations (1) and (2); making a steel pipe having a predetermined dimension from the steel pipe raw material; and performing quenching-tempering treatment for the steel pile, in which the

steel pipe is reheated to a temperature of <u>about 850°C</u> or more, is then cooled to <u>about 100°C</u> or less at a cooling rate faster than that of air cooling, and is again heated to a temperature of <u>about 700°C</u> or less, the equations being:

$$Cr+0.65Ni+0.6Mo+0.55Cu-20C \ge 19.5$$
 (1)

$$Cr+Mo+0.3Si-43.5C-0.4Mn-Ni-0.3Cu-9N\geq11.5$$
 (2)

(where Cr, Ni, Mo, Cu, C, Si, Mn, and N represent the respective contents on a mass percent basis).

- (14) According to the above (13), in the method for manufacturing a high strength stainless steel pipe for use in oil wells, pipe making is Pipe making may be performed by hot working while the steel pipe raw material is heated, and cooling is may then be performed to room temperature at a cooling rate faster than that of air cooling so as to form the seamless steel pipe having a predetermined dimension, followed by the above quenching-tempering treatment.
- (15) According to the above (13) or (14), in the method for manufacturing a high strength stainless steel pipe for use in oil wells, instead Instead of the above quenching-tempering treatment, tempering treatment is may be performed by heating the steel pipe to a temperature of above 700°C or less.
- high strength stainless steel pipe for use in oil wells, in In addition to the above composition in the method, the steel pipe raw material may further contains about 0.002% to about 0.05% of Al on a mass percent basis.

- (17) According to one of the above (13) to (16), in In the method for manufacturing a high strength stainless steel-pipe for use in oil wells, the content of C is may be in the range of about 0.03% to about 0.05%.
- (18) According to one of the above (13) to (17), in In the method-for manufacturing a high strength stainless steel pipe for use in oil wells, the content of Cr is may be in the range of about 16.6% to less than about 18%.
- (19) According to one of the above (13) to (18), in In the method for manufacturing a high strength stainless steel pipe for use in oil wells, the content of Mo is may be in the range of about 2% to about 3.5% on a mass percent basis.
- (20) According to one of the above (13) to (19), in In the method for manufacturing a high strength stainless steel pipe for use in oil wells, in addition to the above composition, the steel pipe raw material may further contains about 3.5% or less of Cu on a mass percent basis.
- (21) According to the above (20), in In the method-for manufacturing a high strength stainless steel-pipe for use in oil wells, the content of Cu is may be in the range of about 0.5% to about 1.14% on a mass percent basis.
- According to one of the above (13) to (21), in In the method for manufacturing a high strength stainless steel pipe for use in oil wells, in addition to the above composition, the steel pipe raw material may further contains at least one of about 0.2% or less of Nb, about 0.3% or less of Ti, about 0.2% or less of Zr, about 3% or less of W, and about 0.01% or less of B on a mass percent basis.
- (23) According to one of the above (13) to (22), in In the method for manufacturing a high strength stainless steel pipe for use in oil wells, in addition to the above

composition, the steel pipe raw material <u>may</u> further contains <u>about</u> 0.01% or less of Ca on a mass percent basis.

Brief Description of the Drawings

[0026] Fig. 1 is a graph showing the relationship between the crack length and the value of the left-hand side of equation (2).

[0027] Fig. 2 is a graph showing the relationship between the crack length and the amount of ferrite.

[0028] Fig. 3 is a graph showing the relationship between the corrosion rate and the value of the left-hand side of equation (1).

[0029] Fig. 4 is a graph showing the influence of a texture on the relationship between a yield strength YS and the amount of Cr.

Best Mode for Carrying Out the Invention Detailed Description

In manufacturing a conventional martensite stainless steel seamless pipe, when a martensite single phase is not obtained due to the formation of a ferrite phase, the strength is decreased and hot workability is degraded. Hence, it has been generally believed that manufacturing of the steel pipe cannot be easily performed. Accordingly, as disclosed in Japanese Unexamined Application 8-246107, generally in a 13%Cr stainless steel seamless pipe for use in oil wells, for manufacturing, the composition thereof has been controlled so that the formation of ferrite is suppressed to obtain a texture formed of a martensite single phase.

We found that, when the steel composition is controlled to satisfy the following equation (2), the hot workability is significantly improved, and generation of cracks in hot working can be prevented:

$Cr+Mo+0.3Si-43.5C-0.4Mn-Ni-0.3Cu-9N \ge 11.5$ (2)

(where Cr, Ni, Mo, Cu, C, Si, Mn, and N represent respective contents on a mass percent basis).

Fig. 1 shows the relationship between the value of the left-hand side of the equation [0032] (2) and the length of crack generated in an end surface of a 13%Cr stainless steel seamless pipe in hot working (that is, in pipe making of a seamless steel pipe). As can be seen from Fig. 1, it is understood that, when the value of the left-hand side of the equation (2) is 8.0 or less, or the lefthand side of the equation (2) is 11.5 or more and is preferably 12.0 or more, the generation of cracks can be prevented. A value of the left-hand side of the equation (2) of 8.0 or less represents a region in which ferrite is not formed at all, and this region corresponds to a region defined by the conventional concept of improvement in hot workability in which the formation of a ferrite phase is not allowed. In addition, as the value of the left-hand side of the equation (2) is increased, the amount of ferrite thus formed is increased, and in the region in which the value of the left-hand side is 11.5 or more, a relatively large amount of ferrite is formed. That is, we found that when the concept is employed that is totally different from the conventional one in the past, that is, when the composition is adjusted to have a value of the left-hand side of 11.5 or more so that a texture containing a relatively large amount of ferrite is used in pipe making, the hot workability can be significantly improved.

Fig. 2 shows the relationship between the amount of ferrite and the length of crack generated in the end surface of a 13%Cr stainless steel seamless pipe in hot working, the relationship being obtained based on the data described above. As can be seen from Fig. 2, as is

the conventional concept, cracks are not generated when the amount of ferrite is 0 percent by volume. However, as ferrite is formed, cracking starts to occur. When the amount of ferrite is further increased to 10 percent by volume or more and preferably 15 percent by volume or more, generation of cracks can be prevented, and this phenomenon is totally different from that based on the conventional concept. That is, when the components are adjusted to satisfy the equation (2), and a ferrite-martensite dual phase is formed in which an appropriate amount of a ferrite phase is formed, the hot workability is improved, and the generation of cracks can be prevented. In addition, it was also found that, when a ferrite-martensite dual phase texture is used, a strength required for oil well pipes can also be ensured.

However, when the components are adjusted to satisfy the equation (2) to form a ferrite-martensite dual phase texture, the corrosion resistance may be degraded in some cases due to the distribution of elements which occurs during heat treatment. When the dual phase texture is formed, since elements such as C, Ni, and Cu forming an austenite phase are diffused to a martensite phase, and elements such as Cr and Mo forming a ferrite phase are diffused to a ferrite phase, as a result, variation between the phases occurs in the final product obtained after heat treatment. In the martensite phase, since the amount of Cr effective for corrosion resistance is decreased, and the amount of C degrading corrosion resistance is increased, as a result, the corrosion resistance may be degraded in some cases as compared to that of a uniform texture.

[0035] We also found that, by adjusting components to satisfy the following equation (1), even when a ferrite-austenite dual phase texture is formed, sufficient corrosion resistance can be ensured:

$$Cr+0.65Ni+0.6Mo+0.55Cu-20C \ge 19.5$$
 (1)

(where Cr, Ni, Mo, Cu, and C represent the respective contents on a mass percent basis).

[0036] Fig. 3 shows the relationship between the value of the left-hand side of the equation (1), even when a ferrite-austenite dual phase texture is formed, in a high temperature environment at 230°C containing CO₂ and Cl⁻, sufficient corrosion resistance can be ensured.

As apparent from equation (1), the content of Cr is advantageously increased to improve the corrosion resistance. However, Cr promotes the formation of ferrite. Hence, in order to suppress the formation of ferrite, Ni in an amount corresponding to the content of Cr was necessary to be added in the past. However, when the content of Ni is increased to correspond to the content of Cr, an austenite phase is stabilized and, as a result, a problem may arise in that the strength required for oil well pipes cannot be ensured.

We found that, when the content of Cr is increased while a ferrite-austenite dual phase texture containing an appropriate amount of a ferrite phase is maintained, a remaining amount of an austenite phase can be reduced and a sufficient strength as an oil well pipe can be ensured.

Fig. 4 shows the relationship between the content of Cr and the yield strength YS of a 13%Cr stainless steel seamless pipe containing a ferrite-austenite dual phase texture processed by heat treatment. In Fig. 4, the relationship[between the content of Cr and the yield strength YS of a martensite single phase texture or a martensite-austenite dual phase texture processed by heat treatment is also shown. From Fig. 4, it was first found that when the ferrite-austenite dual phase texture containing an appropriate amount of a ferrite phase is maintained, and the content of Cr is increased, a sufficient strength as an oil well pipe can be ensured. On the other hand, when the texture is a martensite single phase or a martensite-austenite dual phase texture, as the amount of Cr is increased, the YS is decreased.

[0040] First, the The reason the composition of the high strength stainless steel pipe for use in oil wells is restricted in a specific range will be described below. Hereinafter, the content on a mass percent basis will be simply represented by %.

C: about 0.005% or more to about 0.05% or less

[0041] C is an important element relating to the strength of martensite stainless steel and is required to have a content of about 0.005% or more; however. However, when the content is more than about 0.05%, the degree of sensitization in tempering caused by contained Ni is increased. In order to prevent this sensitization, the The content of C is set in the range of about 0.005% to about 0.05% in the present invention to prevent this sensitization. In addition, in view of corrosion resistance, a smaller amount of C is more preferable; however. However, in order to ensure the strength, a large amount of C is preferable. In consideration of the balance therebetween, the content of C is preferably in the range of about 0.03% to about 0.05%.

Si: about 0.05% or more to about 0.5% or less

[0042] Si is an element functioning as a deoxidizing agent, and <u>about 0.05</u>% or more of Si is contained—in the present invention. However, when the content is more than <u>about 0.5</u>%, CO₂ corrosion resistance is degraded, and in addition, the hot workability is also degraded. Hence, the content of Si is set in the range of <u>about 0.05</u>% to <u>about 0.5</u>%. In addition, the content is preferably in the range of <u>about 0.1</u>% to <u>about 0.3</u>%.

Mn: <u>about 0.2%</u> or more to <u>about 1.8%</u> or less

[0043] Mn is an element increasing the strength, and in order to ensure a desired strength in the present invention, the content of Mn is required to beabout 0.2% or more; however. However, when the content is more than about 1.8%, the toughness is adversely influenced. Hence, the content of Mn is set in the range of about 0.2% to about 1.8%. In addition, the

content is preferably in the range of <u>about 0.2%</u> to <u>about 1.0%</u> and more preferably in the range of <u>about 0.2%</u> to <u>about 0.8%</u>.

P: about 0.03% or less

[0044] P is an element degrading the CO₂ corrosion resistance, resistance to CO₂ stress corrosion cracking, pitting resistance, and resistance to sulfide stress cracking, and hence the content of P is preferably decreased as small as possible in the present invention; however. However, when the content is excessively decreased, the manufacturing cost is inevitably increased. As the content which can be obtained at an inexpensive cost from an industrial point of view and which may not degrade the CO₂ corrosion resistance, resistance to CO₂ stress corrosion cracking, pitting resistance, and resistance to sulfide stress cracking, the content of P is set to about 0.03% or less. In addition, the content is preferably about 0.02% or less.

S: about 0.005% or less

[0045] S is an element seriously degrading the hot workability in a pipe manufacturing process, and hence the content thereof is preferably decreased as small as possible. However, when the content is decreased to <u>about 0.005</u>% or less, since pipe manufacturing can be performed by using a common process, the content of S is set to <u>about 0.005</u>% or less. In addition, the content is preferably <u>about 0.002</u>% or less.

Cr: <u>about 15.5%</u> or more to <u>about 18%</u> or less

[0046] Cr is an element improving the corrosion resistance by forming a protective film and, in particular, is an element improving the CO₂ corrosion resistance and the resistance to CO₂ stress corrosion cracking. In order to To improve the corrosion resistance at a high temperature, in particular, the content is required to be about 15.5% or more in the present invention. On the other hand, when the content is more than about 18%, the hot workability is degraded, and, in

addition, the strength is also decreased decreases. Hence, in the present invention, the content of Cr is set in the range of about 15.5% to about 18%. In addition, the content is preferably in the range of about 16.5% to about 18% and more preferably in the range of about 16.6% to less than about 18%.

Ni: about 1.5% or more to about 5% or less

[0047] Ni has—functions to make the protective film stronger and to—improve the CO₂ corrosion resistance, resistance to CO₂ stress corrosion cracking, pitting resistance, and resistance to sulfide stress cracking. The above functions can be obtained when the content is about 1.5% or more; however. However, when the content is more than about 5%, the stability of the martensite texture is degraded, and the strength is decreased. Hence, the content of Ni is set in the range of about 1.5% to about 5%. In addition, the content is preferably in the range of about 2.5% to about 4.5%.

Mo: about 1% or more to about 3.5% or less

[0048] Mo is an element increasing the resistance to pitting corrosion caused by Cl⁻, and in the present invention, the content of Mo is required to be about 1% or more. When the content is less than about 1%, the corrosion resistance is not sufficient in a severe corrosive environment at a high temperature. On the other hand, when the content is more than about 3.5%, in addition to the decrease in strength, the cost is increased. Hence, the content of Mo is set in the range of about 1% to about 3.5%. In addition, the content is preferably in the range of more than about 2% to about 3.5%.

V: about 0.02% or more to about 0.2% or less

[0049] V has effects to increase the strength and to-improve the resistance to stress corrosion cracking. The effects as described above become significant when the content is about 0.02% or

more; however. However, when the content is more than <u>about 0.2%</u>, the toughness is degraded. Hence, the content of V is set in the range of <u>about 0.02%</u> to <u>about 0.2%</u>. In addition, the content is preferably in the range of <u>about 0.02%</u> to <u>about 0.08%</u>.

N: about 0.01% or more to about 0.15% or less

[0050] N is an element improving the pitting resistance, and the content thereof is set to about 0.01% or more—in—the present invention; however. However, when the content is more than about 0.15%, various nitrides are formed, and as a result, the toughness is degraded. Hence, the content of N is set in the range of about 0.01% to about 0.15%. In addition, the content is preferably in the range of about 0.02% to about 0.08%.

O: about 0.006% or less

[0051] O is present in the form of oxides in steel and has adverse influences on various properties; hence. Hence, the content of O is preferably decreased as small as possible for improving the properties. In particular, when the content of O is more than about 0.006%, the hot workability, resistance to CO₂ stress corrosion cracking, pitting resistance, resistance to sulfide stress cracking, and toughness are seriously degraded. Hence, in the present invention, the content of O is set to about 0.006% or less.

[0052] In addition to the above basic composition, in the present invention, about 0.002% to about 0.05% of Al may also be contained. Al is an element having a strong deoxidizing effect, and in order to obtain the above effect, the content is preferably about 0.002% or more; however. However, when the content is more than about 0.05%, the toughenstoughness is adversely influenced. Hence, when Al is contained, the content thereof is preferably set in the range of about 0.002% to about 0.05%. In addition, the content is more preferably about 0.03% or less. When Al is not contained, Al in a content of approximately less than about 0.002% is allowable

as an unavoidable impurity. When the content of Al is controlled to approximately less than about 0.002%, an advantage in that low temperature toughness is significantly increased can be obtained.

[0053] In addition to the above components described above, <u>about 3.5%</u> or less of Cu may be further contained in the present invention. Cu is an element which makes the protective film strong, prevents hydrogen from penetrating steel, and improves the resistance to sulfide stress cracking, and when the content is <u>about 0.5%</u> or more, the above effects become significant. However, when the content is more than <u>about 3.5%</u>, grain boundary precipitation of CuS occurs, and as a result, the hot workability is degraded. Hence, the content of Cu is preferably set to <u>about 3.5%</u> or less. In addition, the content is more preferably in the range of <u>about 0.8%</u> to <u>about 2.5%</u> and even more preferably in the range of <u>about 0.5%</u> to <u>about 1.14%</u>.

[0054] In the present invention, in addition to the components described above, at least one element selected from about 0.2% or less of Nb, about 0.3% or less of Ti, about 0.2% or less of Zr, about 3% or less of W, and about 0.01% or less of B may be further contained.

[0055] Nb, Ti, Zr, W, and B are elements each increasing the strength and may be selectively contained whenever necessary. In addition, Ti, Zr, W, and B are also elements improving the resistance to stress corrosion cracking. The effects described above become significant, when about 0.03% or more of Nb, about 0.03% or more of Ti, about 0.03% or more of Zr, about 0.2% or more of W, or about 0.0005% or more of B is contained. On the other hand, when more than about 0.2% of Nb, more than about 0.3% of Ti, more than about 0.2% of Zr, more than about 3% of W, or more than about 0.01% of B is contained, the toughness is degraded. Hence, the contents of Nb, Ti, Zr, W, and B are preferably set to about 0.2% or less, about 0.3% or less, about 0.2% or less, about 0.3% or less, and about 0.01% or less, respectively.

[0056] In addition to the above components described above, in the present invention, about 0.01% or less of Ca may also be contained. Ca fixes S by forming CaS and serves to spheroidize sulfide inclusions; hence. Hence, lattice strains of matrix in the vicinity of the inclusions are decreased, so that an effect of decreasing hydrogen trapping ability of the inclusions can be obtained. The effect described above becomes significant when the content is about 0.0005% or more; however. However, when the content is more than about 0.01%, the amount of CaO is increased, and as a result, the CO₂ corrosion resistance and the pitting resistance are degraded. Hence, the content of Ca is preferably set to about 0.01% or less.

[0057] In the present invention, while While being within the ranges described above, the contents of the above components are adjusted so as to satisfy the following equations (1) and (2):

$$Cr+0.65Ni+0.6Mo+0.55Cu-20C \ge 19.5$$
 (1)

$$Cr+Mo+0.3Si-43.5C-0.4Mn-Ni-0.3Cu-9N\ge11.5$$
 (2).

[0058] In the above equations, Cr, Ni, Mo, Cu, C, Si, Mn, and N represent the respective contents (percent by mass). In addition, when the left-hand sides of the equations (1) and (2) are calculated, the content of an element which is not contained is regarded as 0% for calculation.

[0059] When the contents of Cr, Ni, Mo, Cu, and C are adjusted so as to satisfy the equation (1), corrosion resistance in a corrosive environment in which the temperature is high, such as up to 230°C, and CO₂ and Cl⁻ are present can be significantly improved. In addition, in view of improvement in corrosion resistance in a high temperature corrosive environment containing CO₂ and Cl⁻, the value of the left-hand side of the equation (1) is preferably set to 20.0 or more.

[0060] In addition, when the contents of Cr, Mo, Si, C, Mn, Ni, Cu, and N are adjusted to satisfy the equation (2), the hot workability is improved. In the present invention, in order to

improve the hot workability, the The contents of P, S, and O are considerably decreased; however to improve hot workability. However, when the contents of P, S, and O are each only decreased, sufficient and enough hot workability cannot be ensured for making a martensite stainless steel seamless pipe. In order to To ensure sufficient and enough hot workability for making a stainless steel seamless pipe, in addition to a decrease in content of P, S, and O, it is important that the contents of Cr, Mo, Si, C, Mn, Ni, Cu, and N are adjusted to satisfy the equation (2). In addition, in view of improvement in hot workability, the value of the left-hand side of the equation (2) is preferably set to 12.0 or more.

[0061] The balance other than the components described above includes Fe and unavoidable impurities.

[0062] In addition to the components described above, the high strength stainless steel pipe for use in oil wells, according to the present invention, preferably has a texture containing a martensite phase as a primary phase and a ferrite phase at a volume fraction of about 10% to about 60% and preferably of more than about 10% to about 60%.

[0063] In order to ensure a high strength, the The steel pipe of the present invention contains a martensite texture as a primary texture to ensure high strength. In order to improve the toughness without decreasing the strength, the The texture preferably contains a martensite phase as a primary phase and a ferrite phase as a second phase at a volume fraction of about 10% to about 60% and preferably of more than about 10% to about 60% to improve the toughness without decreasing the strength. When the ferrite phase is less than 10 percent by volume or about 10 percent by volume or less, a predetermined object cannot be achieved. On the other hand, when more than about 60 percent by volume of the ferrite phase is contained, the strength is decreased. Hence, the volume fraction of the ferrite phase is set in the range of about 10% to

<u>about 60%</u> and is preferably set in the range of more than <u>about 10%</u> to <u>about 60%</u>. In addition, more preferably, the volume fraction is in the range of <u>about 15%</u> to <u>about 50%</u>. As the second phase other than the ferrite phase, when an austenite phase at a volume fraction of <u>about 30%</u> or less is contained, no problems may arise at all.

[0064] Next, a method for manufacturing a steel pipe, according to the present invention, will be described using a seamless steel pipe by way of example.

[0065] It is preferable that, first, molten steel having the composition described above is formed into an ingot by a known ingot-forming method using a converter, an electric furnace, a vacuum melting furnace, or the like, followed by formation of steel pipe raw materials such as billets using a known method including a continuous casting method or an ingot making-bloom rolling method. Next, these steel pipe raw materials are heated and processed by hot working for making a pipe using a manufacturing process such as a general Mannesmann-plug mill method or Mannesmann-mandrel mill method, so that a seamless steel pipe having a desired dimension is formed. After the pipe makingpipe making, the seamless steel pipe is preferably cooled to room temperature at a cooling rate faster than that of air cooling. Alternatively, the seamless steel pipe may be manufactured by hot extrusion using a press method.

[0066] When a seamless steel pipe has the <u>above described</u> composition within the range of the present invention, a texture having a martensite phase as a primary phase can be formed by hot working, followed by cooling to room temperature at a cooling rate faster than that of air cooling. However, it is preferable that, after the <u>pipe makingpipe making</u> and following the cooling at a cooling rate faster than that of air cooling, quenching treatment be performed in which reheating is performed to a temperature of <u>about 850°C</u> or more, followed by cooling to <u>about 100°C</u> or less and preferably to room temperature at a cooling rate faster than that of air

cooling. By the above treatment, preferably, a fine and tough martensite texture containing an appropriate amount of a ferrite phase can be obtained.

[0067] When the quenching temperature is less than <u>about</u> 850°C, sufficient quenching cannot be performed for a martensite portion, and as a result, the strength tends to decrease. Hence, the heating temperature in the quenching treatment is preferably set to <u>about</u> 850°C or more.

Subsequently, the seamless steel pipe processed by the quenching treatment is preferably processed by tempering treatment in which the steel pipe is heated to a temperature of about 700°C or less, followed by cooling at a cooling rate faster than that of air cooling. By tempering treatment in which heating is performed to about 700°C or less and preferably to about 400°C or more, a texture is obtained which is formed of a tempered martensite phase or is formed of the tempered martensite phase together with small amounts of a ferrite phase and an austenite phase, so that a seamless steel pipe can be obtained having a desired high toughness and desired superior corrosion resistance besides a desired high strength.

[0069] Alternatively, the tempering treatment may only be performed without performing the quenching treatment.

[0070] The presentSelected aspects of the invention hashave been described using the seamless steel pipe by way of example; however, the present invention is. However, those aspects are not limited thereto. By using a steel pipe raw material having the composition within the above described range of the present invention, and in accordance with a common manufacturing process, an electric resistance welded steel pipe and a UOE steel pipe can be manufactured as an oil-well steel pipe.

[0071] For steel pipes other than the seamless steel pipe, such as an electric resistance welded steel pipe and a UOE steel pipe, which are obtained in accordance with a common manufacturing process using a steel pipe raw material having the composition within the range of the present inventiondescribed above, the quenching-tempering treatment described above is preferably performed after pipe-makingpipe making. That is, it is preferable that the quenching treatment be performed in which reheating is performed to a temperature of about 850°C or more, followed by cooling to about 100°C or less and preferably to room temperature at a cooling rate faster than that of air cooling, and that the tempering treatment be then performed in which heating is performed to about 700°C or less and preferably to about 400°C or more, followed by cooling at a cooling rate faster than that of air cooling.

Examples

[0072] Next, selected aspects of the present-invention will be further described in detail with reference to the examples.

Example 1

[0073] After degassing was performed, molten steel having the composition shown in Table 1 was cast into a steel ingot (steel pipe raw material) in an amount of 100 kg, followed by hot working using a model seamless rolling mill for pipe-makingpipe making. After the pipe-makingpipe making, air cooling or water cooling was performed, so that a seamless steel pipe (having an outer diameter of \$3883.8 mm and a wall thickness of 12.7 mm (3.3 inches and 0.5 inches in wall thickness) was obtained.

[0074] The seamless steel pipe thus obtained was examined by visual inspection whether cracks were generated in the inner and the outer surfaces while the steel pipe was placed in a state of air cooling performed after the pipe-makingpipe making, so that the hot workability was

evaluated. When a crack having a length of 5 mm or more was present in the front and the rear

end surfaces of the pipe, it was determined that a crack was generated, and in the other cases, it

was determined that no cracks were generated.

[0075] In addition, from the seamless steel pipe thus obtained, a test piece raw material was

formed by cutting and was heated to 920°C for 30 minutes, followed by water cooling (800%°C)

or more, at an average cooling rate of 10°C/second to 500°C). Furthermore, tempering treatment

at 580°C for 30 minutes was performed. A test piece for texture observation was obtained from

the test piece raw material processed by the above quenching-tempering treatment, followed by

corrosion treatment using aqua regia. Subsequently, an image of the texture of the test piece was

taken using a scanning electron microscope (at 1,000 magnifications), and by using an image

analysis device, the fraction (percent by volume) of a ferrite phase was calculated.

[0076] In addition, the fraction of a retained austenite phase was also measured by an x-ray

diffraction method. After a test piece for measurement was obtained from the test piece raw

material processed by the quenching-tempering treatment, the diffracted x-ray integrated

intensity of the (220) plane of γ and that of the (211) plane of α were measured using an x-ray

diffraction method and were then converted by the following equation. By the way, the fraction

of the martensite phase was calculated as a remaining part other than the phases described above.

 γ (volume fraction) = $100/\{1+(I\alpha R\gamma/I\gamma R\alpha)\}$

[0077] In the above equation, the symbols are:

Ia: integrated intensity of α ,

Iy: integrated intensity of γ ,

R α : crystallographic theoretical calculation value of α ,

Ry: crystallographic theoretical calculation value of γ .

29

[0078] In addition, after an arc-shaped API tensile test piece was formed from the test piece raw material processed by the quenching-tempering treatment, a tensile test was performed, so that the tensile properties (yield strength YS and tensile strength TS) were obtained.

[0079] Furthermore, a corrosion test piece having a thickness of 3 mm, a width of 30 mm, and a length of 40 mm was formed by machining from the test piece raw material processed by the quenching-tempering treatment, and a corrosion test was then performed.

[0080] In the corrosion test, the corrosion test piece was immersed in an aqueous test solution containing 20% of NaCl (at a solution temperature of 230°C under 100 atmospheric pressure in a CO₂ gas atmosphere) placed in an autoclave and was held for 2 weeks as an immersion period. The weight of the corrosion test piece after the corrosion test was measured, and from the reduction in weight before and after the corrosion test, the corrosion rate was obtained by calculation. In addition, by using the corrosion test piece after the corrosion test, the presence of pitting generated in the surface of the test piece was observed using a loupe having a magnification of 10x. When a pitting hole having a diameter of 0.2 mm or more was formed by pitting, it was determined that pitting occurred, and in the other cases, it was determined that no pitting occurred. The results are shown in Table 2.

Table 1

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Remarks		Example	Example	Example	Example	Example	Example	Example	Example	Comparative example	Example	Comparative example	Example	Example	Example	Example	Example	Example	Example	Example				
Value of left-hand	side of equation (2)**	13.19	14.64	14.40	14.97	12.91	14.57	14.39	13.26	14.88	11.73	11.24	11.33	12.49	15.59	17.42	16.37	17.76	13.73	14.97	13.24	13.36	14.11	14.35
Value of left-hand	side of equation (1)*	20.04	20.85	20.40	20.29	20.57	20.76	21.59	20.28	19.29	19.57	19.87	20.36	18.97	20.75	19.59	20.88	21.13	19.55	19.66	20.56	21.56	21.77	21.94
	Ca	-	ı	-	•	•	0.002	0.001	•	-	•	-	-	ı	-	•	•	1	ı	ı	-	•	•	•
	Nb, Ti, Zr, W,B	1	Nb:0.068	Ti:0.036	Zr:0.025	Ti:0.021,B:0.001	•	Nb:0.044	W:0.26	•	•	Ti:0.025	•	Nb:0.061	•	•	Nb:0.077	Ti:0.064	1	Nb:0.083	Nb:0.061	Nb:0.056	Ti:0.049	Nb:0.073
	Cu	0.98	,	ı	08.0	1.24	,	0.75	•	•	1.16	0.62	1.18	*	•	•	0.22	1	1	0.15	0.94	86.0	1.05	1.02
nts	0	0.0031	0.0023	0.0270	0.0035	0.0028	0.0025	0.0019	0.0016	0.0028	0.0017	0.0024	0.0026	0.0034	0.0018	0.0030	0.0038	0.0025	0.0031	0.0036	0.0026	0.0033	0.0019	0.0022
Chemical components	Z	0.047	0.053	0.057	0.062	0.044	0.039	0.050	0.063	0.056	0.106	0.042	0.059	0.058	0.045	0.051	0.031	0.024	0.037	0.026	0.044	0.051	0.039	0.028
Chemica	>	0.047	0.057	0.059	0.049	0.038	0.051	0.046	0.055	0.046	0.055	0.048	0.063	0.065	0.056	0.060	0.061	0.052	0.055	0.053	0.052	0.049	0.061	0.053
	AI	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.02	0.01	0.01	0.02	0.02	0.01	0.02	0.02	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
	Mo	1.6	2.5	2.4	2.1	1.9	1.8	2.4	2.0	1.9	2.7	1.9	1.5	0.4	2.6	2.9	2.7	3.3	2.6	2.8	2.6	2.0	3.1	3.2
	ïZ	3.5	3.7	3.6	2.6	3.8	3.6	4.0	3.9	2.6	3.8	4.6	4.6	3.3	2.9	10	2.8	2.5	3.8	3.1	3.7	7.1	4.2	4.0
	ပ်	16.6	17.4	17.0	17.4	16.8	17.8	17.5	17.2	16.7	15.4	16.1	16.3	17.1	17.3	17.2	17.7	17.8	15.7	16.1	16.9	17.9	17.1	17.7
	S	0.002	0.001	0.001	0.001	0.002	0.001	0.001	0.001	0.001	0.001	0.001	0.002	0.001	0.001	0.001	0.001	0.001	0.001	0.001	0.001	0.001	0.001	0.001
	Ь	0.01	0.01	0.01	0.02	0.02	0.02	0.02	0.01	0.02	0.02	0.02	0.01	0.02	0.02	0.01	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.02
	Mn	0.26	0.35	0:30	0.29	0.38	0.36	0.31	0.27	0.45	0.35	0.39	0.35	0.36	0.32	0:30	0.28	0.25	0.31	0.35	0.41	0.39	0.52	0.61
	. <u>S</u>	0.19	0.18	0.21	0.23	0.20	0.21	0.23	0.25	0.27	0.29	0.28	0.24	0.29	0.25	0.26	0.17	0.28	0.25	0.24	0.22	0.25	0.23	0.25
	O	0.017	0.023	0.019	0.025	0.026	0.023	0.018	0.033	0.012	0.028	0.035	0.023	0.026	0.012	0.027	0.019	0.014	0.009	0.011	0.041	0.037	0.025	0.042
Steel No.		∢	മ	ပ	۵	Ш	щ	ပ	I	-		エ		Σ	z	0	۵.	a	α.	S	⊢	כ	>	3

Left-hand side of equation (1): Cr+0.65Ni+0.6Mo+0.55Cu-20C

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Left-hand side of equation (2): Cr+Mo+0.3Si -43.5C-0.4Mn -Ni-0.3Cu -9N

Table 2

9.

(i) Grandk			Hot workability		Com	Composition		Tensile p	properties	Corrosion	Corrosion resistance	
Waler cooling - M+F+Y 75.8 13.5 10.7 823 984 0.108 Ak cooling No M+F+Y 75.2 14.6 12.2 819 990 0.104 Ak cooling No M+F+Y 56.1 30.2 17.9 843 994 0.083 Aki cooling No M+F+Y 56.2 26.7 18.8 838 999 0.101 Aki cooling No M+F+Y 66.4 15.2 17.9 843 994 0.001 Aki cooling No M+F+Y 66.4 15.2 18.4 823 999 0.101 Aki cooling No M+F+Y 66.4 15.2 18.4 823 990 0.105 Aki cooling No M+F+Y 57.9 28.1 18.0 973 0.105 Aki cooling No M+F+Y 78.2 10.2 11.6 75.3 994 0.104 Aki cooling No <	Ste. No.		Presence of crack generation	Types*	Amount of martensite (percent by volume)	Amount of ferrite (percent by volume)	Amount of austenite (percent by volume)		TS (MPa)	Corrosion rate (mm/yr)	Presence of pitting generation	Remarks
Air cooling No MAFFTY 73.2 14.6 12.2 819 980 0.114 Air cooling No MAFFTY 55.1 30.3 14.6 684 996 0.013 Water cooling No MAFFTY 56.9 25.2 17.9 684 996 0.010 Air cooling No MAFTY 65.4 15.2 18.8 687 0.097 Air cooling No MAFTY 65.4 15.2 19.4 62.3 980 0.105 Air cooling No MAFTY 65.4 15.2 19.4 62.3 980 0.105 Air cooling No MAFTY 66.9 17.4 15.2 984 0.016 Air cooling No MAFTY 66.9 17.4 15.2 984 981 0.105 Air cooling No MAFTY 78.2 28.1 16.2 97.2 0.105 Air cooling No MAFTY 78.6 <td><</td> <td>Water cooling</td> <td>•</td> <td>$M+F+\gamma$</td> <td>75.8</td> <td>13.5</td> <td>10.7</td> <td>823</td> <td>984</td> <td>0.108</td> <td>No</td> <td>Example</td>	<	Water cooling	•	$M+F+\gamma$	75.8	13.5	10.7	823	984	0.108	No	Example
Aut cooling No MHF+y \$5.1 30.3 14.6 964 996 0.087 Water cooling MHF+y 56.3 28.2 17.9 84.3 994 0.037 Air cooling No MHF+y 62.3 22.9 4.8 88.7 1008 0.101 Air cooling No MHF+y 65.4 15.2 19.4 82.3 980 0.105 Air cooling No MHF+y 57.8 28.4 13.0 775 974 0.084 Air cooling No MHF+y 57.8 28.4 13.0 775 974 0.084 Air cooling No MHF+y 57.8 28.4 15.7 836 969 0.104 Air cooling No MHF+y 57.8 28.4 6.2 984 0.105 Air cooling No MHF+y 78.2 10.2 11.6 77.5 986 0.104 Air cooling No	∢ .	Air cooling	o Z	M+F+y	73.2	14.6	12.2	819	086	0.114	No	Example
Waler cooling - M+F*y 56.9 25.2 17.9 84.3 994 0.097 Air cooling No M+F*y 54.5 26.7 18.8 888 969 0.101 Air cooling No M+F*y 65.4 15.2 18.4 857 1009 0.105 Air cooling No M+F*y 66.4 15.2 18.4 827 1009 0.105 Air cooling No M+F*y 66.9 28.4 13.0 775 97.4 0.064 Air cooling No M+F*y 66.9 28.4 18.0 981 0.004 Air cooling No M+F*y 78.2 17.4 16.7 889 981 0.104 Air cooling No M+F*y 77.1 1.5 21.4 818 97.2 0.142 Air cooling No M+F*y 78.6 2.8 8.2 980 0.132 Air cooling No M+F*y 78.6	8	Air cooling	No	M+F+y	55.1	30.3	14.6	864	966	0.093	S _O	Example
Africooling No M+F+y 54.5 28.7 18.8 838 989 0.101 Africooling No M+F+y 62.3 32.9 4.8 867 1009 0.105 Africooling No M+F+y 65.4 15.2 19.4 823 960 0.036 Africooling No M+F+y 58.6 28.4 13.0 775 974 0.094 Africooling No M+F+y 66.9 17.4 15.7 836 981 0.016 Africooling No M+F+y 61.4 22.4 62.2 816 97.2 0.142 Africooling No M+F+y 78.2 10.2 11.6 889 981 0.104 Africooling No M+F+y 78.2 28.4 15.2 812 960 0.104 Africooling No M+F+y 77.6 18.1 82.9 842 0.05 Africooling No <	(Water cooling	•	M+F+y	56.9	25.2	17.9	843	994	0.097	No	Example
Air cooling No M+F+y 62.3 32.9 4.8 867 1009 0.105 Air cooling No M+F+y 65.4 15.2 19.4 823 980 0.036 Air cooling No M+F+y 56.9 28.4 13.0 775 974 0.098 Air cooling No M+F+y 66.9 17.4 15.7 836 990 0.104 Air cooling No M+F+y 66.9 17.4 15.7 836 990 0.104 Air cooling No M+F+y 66.9 17.4 15.7 836 990 0.104 Air cooling No M+F+y 78.2 10.2 11.6 872 0.132 Air cooling No M+F+y 78.6 2.9 20.5 829 984 0.086 Air cooling No M+F+y 78.6 18.1 820 981 0.104 Air cooling No M+F+y <	ر	Air cooling	No	Μ+F+γ	54.5	26.7	18.8	838	686	0.101	No	Example
Africooling No M+F+y 65.4 15.2 19.4 823 980 0.098 Air cooling No M+F+y 58.6 28.4 13.0 775 974 0.094 Air cooling No M+F+y 65.9 17.4 15.7 836 981 0.076 Air cooling No M+F+y 65.9 17.4 6.2 849 981 0.046 Air cooling No M+F+y 78.2 10.2 11.6 783 989 0.142 Air cooling No M+F+y 77.1 1.5 21.4 818 97.2 0.142 Air cooling Ves M+F+y 78.6 2.9 20.5 812 989 0.132 Air cooling No M+F+y 78.6 2.9 20.5 82 984 981 0.105 Air cooling No M+F+y 76.6 2.9 20.5 62.9 6.05 0.05 Air coo	D	Air cooling	No	M+F+y	62.3	32.9	4.8	867	1009	0.105	N _O	Example
Air cooling No MHF hy 58.6 28.4 13.0 775 974 0.094 Air cooling No MHF hy 57.9 26.1 16.0 849 981 0.006 Air cooling No MHF hy 66.9 17.4 15.7 836 969 0.104 Air cooling No MHF hy 78.2 10.2 11.6 763 989 0.104 Air cooling No MHF hy 77.1 1.5 21.4 816 972 0.142 Air cooling No MHF hy 76.6 2.9 20.5 812 989 0.132 Air cooling No MHF hy 74.6 16.1 9.3 824 0.05 0.105 Air cooling No MHF hy 74.6 57.2 0 57.3 980 0.105 Air cooling No MHF hy 74.8 57.2 0 68 87 984 0.051 Air	ш	Air cooling	ON	Μ+F+γ	65.4	15.2	19.4	823	980	0.098	No	Example
Air cooling No M+F+y 57.9 26.1 16.0 649 981 0.076 Air cooling No M+F+y 66.9 17.4 15.7 636 969 0.104 Air cooling No M+F+y 61.4 32.4 6.2 816 972 0.104 Air cooling No M+F+y 77.1 1.5 21.4 818 973 0.105 Air cooling Ves M+F+y 76.6 2.9 20.5 812 989 0.134 Air cooling No M+F+y 76.6 2.9 20.5 872 984 0.105 Water cooling No M+F+y 74.6 16.1 9.3 823 984 0.096 Water cooling No M+F+y 46.2 50.9 2.9 681 875 0.097 Air cooling No M+F+y 46.2 50.9 2.9 694 0.075 Air cooling No <td< td=""><td>L</td><td>Air cooling</td><td>oN</td><td>M+F+y</td><td>58.6</td><td>28.4</td><td>13.0</td><td>277</td><td>974</td><td>0.094</td><td>No</td><td>Example</td></td<>	L	Air cooling	oN	M+F+y	58.6	28.4	13.0	277	974	0.094	No	Example
Air cooling No M+F+y 66.9 17.4 15.7 836 969 0.104 Air cooling No M+F+y 61.4 32.4 6.2 816 972 0.102 Air cooling No M+F+y 77.1 1.5 21.4 818 973 0.139 Air cooling Yes M+F+y 77.1 1.5 21.4 818 973 0.135 Air cooling Yes M+F+y 76.6 2.9 20.5 812 989 0.132 Air cooling No M+F+y 74.6 16.1 9.3 824 969 0.134 Air cooling No M+F+y 46.2 50.9 2.9 681 820 0.096 Air cooling No M+F+y 46.2 50.9 2.9 691 0.134 Air cooling No M+F+y 34.5 62.9 2.6 689 875 0.097 Air cooling No M+F	9	Air cooling	ON	M+F+y	67.9	26.1	16.0	849	981	0.076	ON ON	Example
Air cooling No M+F+Y 61.4 32.4 6.2 816 972 0.142 Air cooling No M+F+Y 78.2 10.2 11.6 763 989 0.139 Air cooling Yes M+F+Y 77.1 1.5 21.4 818 973 0.105 Air cooling Yes M+F+Y 76.6 2.9 20.5 812 958 0.132 Air cooling No M+F+Y 76.6 2.9 20.5 82 829 884 0.036 Air cooling No M+F+Y 76.6 33.6 6.8 829 894 0.096 Air cooling No M+F+Y 46.2 57.2 0 57.3 916 0.134 Air cooling No M+F+Y 46.2 50.9 2.9 691 875 0.097 Air cooling No M+F+Y 72.9 27.7 0 100.4 110.5 0.125 Air coolin	エ		No	M+F+y	6.99	17.4	15.7	836	696	0.104	No	Example
Air cooling No M+F+y 78.2 10.2 11.6 76.3 989 0.139 Air cooling Yes M+F+y 77.1 1.5 21.4 818 973 0.105 Air cooling Yes M+F+y 76.6 2.9 20.5 812 958 0.132 Air cooling No M+F+y 76.6 2.9 20.5 829 984 0.105 Water cooling No M+F+y 58.6 33.6 6.8 8.3 821 980 0.174 Air cooling No M+F+y 41.9 57.2 0 57.3 916 0.105 Air cooling No M+F+y 46.2 50.9 2.9 691 87.3 0.081 Air cooling No M+F+y 71.8 28.2 0 107.2 11.9 Air cooling No M+F+y 62.9 27.1 0 10.2 0.095 Air cooling No M+		Air cooling	°N	M+F+y	61.4	32.4	6.2	816	972	0.142	No	Comparative example
Air cooling Yes M+F+y 77.1 1.5 21.4 818 97.3 0.105 Air cooling Yes M+F+y 76.6 2.9 20.5 812 958 0.132 Air cooling No M+F+y 74.6 16.1 9.3 834 969 0.174 Water cooling No M+F+y 59.6 33.6 6.8 829 984 0.096 Water cooling No M+F+y 41.9 57.2 0 6.8 829 0.174 Air cooling No M+F+y 44.9 50.9 2.9 691 820 0.105 Air cooling No M+F+y 72.9 27.1 0 1012 114 0.115 Air cooling No M+F+y 62.7 18.8 18.5 990 0.095 Air cooling No M+F+y 62.7 18.5 870 0.095 0.095 Air cooling No M+F+y <	ال	Air cooling	No	M+F+y	78.2	10.2	11.6	763	989	0.139	No	Comparative example
Air cooling Yes M+F+y 76.6 2.9 20.5 812 958 0.132 Air cooling No M+F+y 74.6 16.1 9.3 834 969 0.174 0.036 Water cooling - M+F+y 59.6 33.6 6.8 829 984 0.096 0.174 Air cooling No M+F+y 41.9 57.2 0 573 916 0.100 0.096 Air cooling No M+F+y 46.2 50.9 2.9 691 892 0.097 0.037 Air cooling No M+F+y 46.2 50.9 2.9 694 1051 0.037 Air cooling No M+F+ 71.8 27.1 0 1012 1114 0.119 Air cooling No M+F+y 62.7 18.8 855 990 0.095 Air cooling No M+F+y 64.3 19.5 16.2 870 0.095 0.	지	Air cooling	Yes	M+F+γ	77.1	1.5	21.4	818	973	0.105	N _O	Comparative example
Air cooling No M+F+y 74.6 16.1 9.3 834 969 0.174 P Water cooling - M+F+y 59.6 33.6 6.8 8.3 829 984 0.096 0.006 Air cooling - M+F+y 41.9 57.2 0 57.3 96 0.006 0.006 Air cooling No M+F+y 46.2 50.9 2.6 669 875 0.081 0.037 Air cooling No M+F+y 34.5 62.9 2.6 669 875 0.081 Air cooling No M+F+y 72.9 27.1 0 1012 1114 0.119 Air cooling No M+F+y 62.7 18.8 18.5 855 990 0.097 1 Air cooling No M+F+y 62.7 18.8 18.5 870 0.095 1 Air cooling No M+F+y 63.7 27.7 18.6	_1	Air cooling	Yes	M+F+y	9.92	2.9	20.5	812	958	0.132	No	Comparative example
Water cooling - M+F+y 59.6 33.6 6.8 829 984 0.096 Air cooling No M+F+y 57.8 33.9 8.3 8.21 980 0.100 Water cooling - M+F+y 41.9 57.2 0 57.3 916 0.134 0 Air cooling No M+F+y 34.5 62.9 2.9 691 875 0.037 0 Air cooling No M+F+y 34.5 62.9 2.6 669 875 0.037 0 Water cooling No M+F 72.9 27.1 0 1012 114 0.119 0 Air cooling No M+F+y 62.7 18.8 18.5 855 990 0.095 0 Air cooling No M+F+y 62.7 27.7 18.6 87 964 0.075 0	≥l		ON .	M+F+y	74.6	16.1	9.3	834	696	0.174	N _O	Comparative example
Air cooling No M+F+y 57.8 33.9 8.3 8.3 8.3 98.0 0.100 Water cooling - M+F+y 41.9 57.2 0 57.3 916 0.134 Air cooling No M+F+y 34.5 62.9 2.9 691 892 0.097 Air cooling No M+F+y 34.5 62.9 2.6 669 875 0.081 Water cooling No M+F 72.9 27.1 0 1012 1114 0.125 Air cooling No M+F+y 62.7 18.8 18.5 855 990 0.097 Air cooling No M+F+y 62.7 18.6 870 1002 0.095 Air cooling No M+F+y 53.7 27.7 18.6 877 959 0.074	Z		•	M+F+y	59.6	33.6	6.8	829	984	0.096	No	Example
Water cooling - M+F+y 41.9 57.2 0 57.3 916 0.134 Air cooling No M+F+y 46.2 50.9 2.9 691 892 0.097 Air cooling No M+F+y 34.5 62.9 2.6 669 875 0.081 Water cooling No M+F 72.9 27.1 0 1012 1114 0.115 Air cooling No M+F+y 62.7 18.8 16.2 0 0 0 0.095 Air cooling No M+F+y 64.3 19.5 16.2 870 0.095 0 Air cooling No M+F+y 52.6 28.1 18.6 837 929 0.074 Air cooling No M+F+y 52.6 28.1 19.3 858 964 0.075	:		No	M+F+y	57.8	33.9	8.3	821	980	0.100	No	Example
Air cooling No M+F+y 46.2 50.9 2.9 691 892 0.097 Air cooling No M+F+y 34.5 62.9 2.6 669 875 0.081 Valer cooling No M+F 72.9 27.1 0 1012 1114 0.125 Air cooling No M+F+y 62.7 18.8 18.5 855 990 0.097 Air cooling No M+F+y 64.3 19.5 16.2 870 0.095 0.095 Air cooling No M+F+y 53.7 27.7 18.6 858 964 0.075	이		•	M+F+y	41.9	57.2	0	573	916	0.134	Yes	Comparative example
Air cooling No M+F+y 34.5 62.9 2.6 669 875 0.081 7 Water cooling No M+F 72.9 27.1 0 1012 1114 0.119 0.119 Air cooling No M+F+y 62.7 18.8 18.5 855 990 0.097 Air cooling No M+F+y 64.3 19.5 16.2 870 1002 0.095 Air cooling No M+F+y 53.7 27.7 18.6 837 929 0.074 Air cooling No M+F+y 52.6 28.1 19.3 858 964 0.075	۵.		o Z	M+F+y	46.2	50.9	2.9	691	892	0.097	0 2	Example
Air cooling No M+F 83.1 16.9 0 964 1051 0.125 7 Water cooling - M+F 72.9 .27.1 0 1012 1114 0.119 7 Air cooling No M+F+y 62.7 18.8 18.5 855 990 0.097 7 Air cooling No M+F+y 64.3 19.5 16.2 870 1002 0.095 7 Air cooling No M+F+y 53.7 27.7 18.6 858 964 0.075 7	g		No	M+F+y	34.5	62.9	2.6	699	875	0.081	No	Example
Water cooling Indeed or cooling <	2		ON.	M+F	83.1	16.9	0	964	1051	0.125	No	Example
Air cooling No M+F+y 71.8 28.2 0 1004 1105 0.122 Air cooling No M+F+y 62.7 18.8 16.2 855 990 0.097 7 Air cooling No M+F+y 53.7 27.7 18.6 837 929 0.074 7 Air cooling No M+F+y 52.6 28.1 19.3 858 964 0.075 7	U.		-	M+F	72.9	. 27.1	0	1012	1114	0.119	No	Example
Air cooling No M+F+γ 62.7 18.8 18.5 855 990 0.097 Air cooling No M+F+γ 53.7 27.7 18.6 837 929 0.074 Air cooling No M+F+γ 52.6 28.1 19.3 858 964 0.075			No	M+F	71.8	28.2	0	1004	1105	0.122	ON	Example
Air cooling No M+F+γ 64.3 19.5 16.2 870 1002 0.095 Air cooling No M+F+γ 53.7 27.7 18.6 837 929 0.074 Air cooling No M+F+γ 52.6 28.1 19.3 858 964 0.075	-	Air cooling	No	M+F+γ	62.7	18.8	18.5	855	066	0.097	N _O	Example
Air cooling No M+F+γ 53.7 27.7 18.6 837 929 0.074 Air cooling No M+F+γ 52.6 28.1 19.3 858 964 0.075)		No	M+F+y	64.3	19.5	16.2	870	1002	0.095	No	Example
Air cooling No $M+F+\gamma$ 52.6 28.1 19.3 858 964 0.075	>		No	M+F+y	53.7	27.7	18.6	837	929	0.074	ON O	Example
	\$		°N	M+F+y	52.6	28.1	19.3	858	964	0.075	o N	Example

M: Martensite, F: Ferrite, γ : Retained austenite

*

[0081] According to examples—of the present invention, the—generation of cracks in the surface of the steel pipe was not observed at all, the yield strength YS was high, such as 654 MPa or more, the corrosion rate was also low, and no pitting occurred; hence. Hence, a steel pipe was obtained having superior hot workability and corrosion resistance in a severe corrosive environment in which CO₂ was present and the temperature was high, such as 230°C. Furthermore, since 5% or more of a ferrite phase was contained, a steel pipe was obtained having high strength, such as a yield strength of 654 MPa or more, and superior corrosion resistance in a severe corrosive environment in which CO₂ was present and the temperature was high, such as 230°C.

On the other hand, according to comparative examples which were outside the range of the present invention, cracks were generated in the surface since the hot workability was degraded; or the corrosion rate was high and pitting occurred since the corrosion resistance was degraded. In particular, in the comparative example in which the equation (2) was not satisfied, the hot workability was degraded, and as a result, scars were generated on the surface of the steel pipe. In addition, when the amount of ferrite was out of the preferable range of the present invention, the strength was decreased, and a high strength, such as a yield strength of 654 MPa or more, could not be achieved.

Example 2

[0083] After the pipe making pipe making was performed by hot working using a steel pipe raw material having the composition (steel No. B, or No. S) shown in Table 1, air cooling was

performed, so that a seamless steel pipe having an outer diameter of 83.8 mm and a wall thickness of 12.7 mm (3.3 inches and 0.5 inches in wall thickness) was obtained. From the seamless steel pipe thus obtained, a test piece raw material was obtained by cutting, followed by quenching-tempering treatment or tempering treatment shown in Table 3.

[0084] A test piece for texture observation and a test piece for measurement were formed from the test piece raw material processed by the quenching-tempering treatment in a manner similar to that in Example 1, and the fraction (percent by volume) of a ferrite phase, the fraction (percent by volume) of a retained austenite phase, and the fraction (percent by volume) of a martensite phase were obtained by calculation.

[0085] In addition, after an arc-shaped API tensile test piece was formed from the test piece raw material processed by the quenching-tempering treatment, a tensile test was performed in a manner similar to that in Example 1, so that the tensile properties (yield strength YS and tensile strength TS) were obtained. Furthermore, in a manner similar to that in Example 1, a corrosion test piece having a thickness of 3 mm, a width of 30 mm, and a length of 40 mm was formed by machining from the test piece raw material processed by the quenching-tempering treatment, and a corrosion test was then performed, so that the corrosion rate was obtained. In addition, in a manner similar to that in Example 1, the presence of pitting generated in the surface of the test piece was observed. The evaluation standard was similar to that in Example 1. The results are shown in Table 3.

Table 3

	1				T .	Ι	Ī			Į.	1				r—	<u> </u>
Remarks		·	Example	Example	Example	Example	Example	Example	Example	Example	Example	Example	Example	Example	Example	Example
Corrosion resistance		Presence of pitting generation	S N	No No	N _o	S	No	No	No	8 N	No	No	No	S	S N	No
Corrosion		Corrosion rate (mm/yr)	0.093	0.101	0.103	660.0	0.095	0.122	0.124	0.122	0.118	960'0	0.094	0.077	960.0	0.097
Tensile properties		TS (MPa)	966	972	955	985	991	1105	1030	1011	1120	995	991	926	986	984
Tensile p		YS (MPa)	864	845	720	850	098	1004	984	896	1014	857	849	805	843	837
		y (percent by volume)	14.6	16.8	21.2	15.1	14.4	0	0	0	0	18.6	18.0	21.9	18.4	17.5
Composition		F (percent by volume)	30.3	32.5	33.0	31.6	30.5	28.2	30.8	29.8	16.8	19.3	18.8	18.6	19.2	17.7
Com		M (percent by volume)	55.1	50.7	45.8	46.7	55.1	71.8	69.2	70.2	73.2	62.1	63.2	59.5	62.4	64.8
		Types*	Μ+F+γ	M+F+y	M+F+y	M+F+y	M+F+y	M+F	¥ ₩	M+F	M+F	M+F+y	M+F+y	M+F+y	M+F+y	M+F+y
	Tempering	Tempering Heating temper- ature (°C)		280	099	085	089	280	650	250	580	580	280	620	280	280
tment	Cooling stop temper- ature (°C)		70	70	70	70	20	02	20	1	70	. 02	20	70	20	70
Heat treatment	Quenching	Cooling	Water cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	•	Air cooling	Air cooling	Air cooling	Air cooling	Water cooling	Air cooling
		Heating temper- ature (°C)	920	920	920	890	860	920	920	ı	890	920	920	920	850	850
	z ciloo	after pipe[[-]]_making	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Water cooling	Air cooling	Air cooling				
	Ste	⊕ °.			മ				ဟ	*				-		
	Steel	pipe No.	2-1	2-2	2-3	2-4	2-5	2-6	2-7	2-8	2-9	2-10	2-11	2-12	2-13	2-14

M: Martensite, F: Ferrite, γ : Retained austenite

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[0086] According to the examples of the present invention, the yield strength YS was high, such as 654 MPa or more, the corrosion rate was also low, and no pitting occurred; hence. Hence, a steel pipe was obtained having superior hot workability and corrosion resistance in a severe corrosive environment in which CO₂ was present and the temperature was high, such as 230°C. However, in examples of the present invention which were out of the preferableour selected range of the present invention, the strength or corrosion resistance and hot workability tend to be degraded.

Example 3

[0087] After degassing was performed, molten steel having the composition shown in Table 4 was cast into an ingot in an amount of 100 kg, followed by hot working using a model seamless rolling mill for pipe makingpipe making. After the pipe makingpipe making, cooling (air cooling) was performed, so that a seamless steel pipe having an outer diameter of 83.8 mm and a wall thickness of 12.7 mm (3.3 inches and 0.5 inches in wall thickness) was obtained.

[0088] The seamless steel pipe thus obtained was examined by visual inspection in a manner similar to that in Example 1 whether cracks were generated in the inner and the outer surface thereof while the steel pipe was placed in a state of air cooling performed after the pipe-making, so that the hot workability was evaluated. In this evaluation, the evaluation standard was similar to that in Example 1.

[0089] In addition, from the seamless steel pipe thus obtained, a test piece raw material was formed by cutting and was heated to 900°C for 30 minutes, followed by water cooling.

Furthermore, tempering treatment at 580°C for 30 minutes was performed. After a test piece for texture observation and a test piece for measurement were obtained from the test piece raw material processed by the quenching-tempering treatment described above, the test piece for texture observation was processed by corrosion treatment using aqua regia. Subsequently, an image of the texture of the test piece was taken using a scanning electron microscope (at 1,000 magnifications), and by an image analysis device, the fraction (percent by volume) of a ferrite phase was calculated. In addition, the test piece for texture observation was obtained from the test piece raw material processed by the quenching-tempering treatment described above, and the fraction (percent by volume) of a retained austenite phase and that of a martensite phase were measured in a manner similar to that in Example 1.

[0090] In addition, after an arc-shaped API tensile test piece was obtained from the test piece raw material processed by the quenching-tempering treatment, a tensile test was performed, so that the tensile properties (yield strength YS and tensile strength TS) were obtained. In addition, after a V notch test piece (thickness: 5 mm) in accordance with JIS Z 2202 was obtained from the test piece raw material processed by the quenching-tempering treatment, a charpy impact test was performed in accordance with JIS Z 2242, so that an absorption energy vE₋₄₀ (J) at -40°C was obtained.

[0091] Furthermore, after a corrosion test piece having a thickness of 3 mm, a width of 30 mm, and a length of 40 mm was formed from the test piece raw material processed by the

quenching-tempering treatment, a corrosion test was performed. By the way, some steel pipe was not processed by the quenching treatment but processed only by the tempering treatment.

[0092] In the corrosion test, the corrosion test piece was immersed in an aqueous test solution containing 20% of NaCl (at a solution temperature of 230°C under 100 atmospheric pressure in a CO₂ gas atmosphere) placed in an autoclave and was held for 2 weeks as an immersion period. The weight of the corrosion test piece after the corrosion test was measured, and from the reduction in weight before and after the corrosion test, the corrosion rate was obtained. In addition, the resistance to pitting was evaluated by immersing the test piece in a solution containing 40% of CaCl₂ (liquid temperature: 70°C) for 24 hours, so that the presence of pitting was examined. When a pitting hole having a diameter of 0.1 mm or more was formed by pitting, it was determined that pitting occurred, and in the other cases, it was determined that no pitting occurred. The results are shown in Table 5.

Table 4

	Remarks	Example	Example	Example	Example	Example	Example	Example	Example	Comparative example	Comparative example	Example	Example	Comparative example	Example	Example	Example	
Value of left-hand	side of equation (2)**	13.34	13.32	13.05	13.72	13.18	14.59	13.43	13.28	13.50	9.88	12.65	15.67	11.48	13.22	13.12	14.09	
Value of left-hand	side of equation (1)*	20.24	20.59	20.34	21.04	20.27	20.89	20.57	21.02	19.35	18.90	20.10	19.80	19.28	21.17	21.43	22.00	
	Al	0.001	0.001	0.001	0.002	0.001	0.001	0.001	0.002	0.001	0.002	0.005	0.012	0.004	0.001	0.001	0.001	
	Ca	-	-		1	-	0.004	0.001	•	•	•	-	0.002	•	ł	•	•	
	Other	•	•	Nb: 0.068	Ti: 0.055	Zr. 0.029 B: 0.001	•	Nb: 0.059	W: 0.48	1	•	Ti: 0.032	ŧ	Nb: 0.038	Nb: 0.061	1	•	
	Cu	•	0.94	,	0.79	1.05	1	0.88	1	1	1.16	0.62	•	•	1.04	0.94	1.11	
	0	0.0029	0.0027	0.0038	0.0044	0.0030	0.0041	0.0038	0.0045	0.0046	0.0038	0.0040	0.0030	0.0026	0.0025	0.0034	0.0020	
y mass)	z	0.050	0.051	0.062	0.058	0.059	0.052	0.042	0.048	0.052	0:050	0.055	0.053	0.051	0.059	0.068	0.055	Cr+0.65Ni+0.6 Mo+0.55Cu-20C Cr+Mo+0.3Si -43.5C-0.4Mn -Ni-0.3Cu -9N
Chemical components (percent by mass)	>	0.049	0.047	0.056	0.059	0.057	0.055	0.047	0.063	0.051	0.047	0.051	0.047	0.060	0.052	0.053	0.002	\u-20C \u-Ni-0
ponents	Mo	1.7	2.4	1.8	1.7	2.1	2.6	2.0	1.7	2.5	1.5	2.1	2.3	1.6	2.2	2.6	3.0	Cr+0.65Ni+0.6 Mo+0.55Cu-20C Cr+Mo+0.3Si -43.5C-0.4Mn -Ni-
cal com	ïZ	4.0	3.8	4.0	3.7	3.4	3.7	3.8	4.4	3.3	4.5	3.9	2.3	4.3	3.9	4.3	4.1	+0.6 M 3Si -43
Chemi	ర	17.0	16.7	17.3	17.7	16.9	17.5	16.8	17.7	16.4	15.0	16.6	17.5	16.2	17.5	17.2	17.6	-0.65Ni -Mo+0.3
	တ	0.001	0.001	0.001	0.001	0.001	0.001	0.002	0.001	0.002	0.001	0.001	0.001	0.001	0.001	0.001	0.001	
	Œ.	0.01	0.02	0.01	0.02	0.02	0.02	0.01	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.02	0.02	equation equation
i	Mn	0.42	0.37	0.45	0.41	0.43	0.39	0.41	0.39	0.42	0.44	0.39	0.40	0.37	0.36	0.42	0.42	Left-hand side of equation (1): Left-hand side of equation (2):
	Si	0.27	0.29	0.28	0.26	0.27	0.26	0.22	0.29	0.20	0.24	0.25	0.24	0.22	0.21	0.26	0.21	Left-han Left-han
	ပ	0.019	0.027	0.032	0.026	0.034	0.029	0.019	0.028	0.035	0.028	0.032	0.029	0.034	0.038	0.032	0.034	£ (**
	No.	14	18	10	10	1后	1F	1G .	1H	1.J	¥	1	1M	Z	1P	10	1R	, T

Table 5

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	Remarks		Example	Example	Example	Example	Example	Example	Example	Example	Comparative example	Comparative example	Example	Example	Comparative example	Example	Example	Example
Pitting resistance	presence	of pitting generation	o _N	o N	92	oN.	°N	^o N	oN.	2	ON.	ON _O	Yes	Yes	Yes	o N	oN N	Š
Corrosion resistance	Corrosion	(mm/y)	0.098	0.094	960'0	0.079	0.098	0.091	0.093	0.088	0.136	0.153	0.102	0.117	0.121	0.095	0.091	0.084
Hot workability	Presence	of crack	oN N	oN N	S _N	ON	8	SN N	No	No	No	Yes	o _N	oN N	ON	No	oN	No
Toughn ess	∨E.40	ſ	91.3	83.5	85.9	87.3	83.3	77.5	87.0	81.7	84.1	99.4	45.4	21.6	46.1	60.5	72.7	85.1
Tensile properties	TS	(MPa)	606	896	963	953	686	915	987	919	933	952	096	906	982	980	696	924
Ter prop	χ	(MPa)	839	826	862	988	228	831	820	668	608	864	842	498	958	828	851	817
ne)	Amount	of ferrite	28.5	31.4	56.6	37.9	30.3	51.2	33.6	31.4	37.3	4.7	22.6	72.3	19.6	19.5	17.6	25.8
ercent by volun	Amount of	retained γ phase	15.2	21.4	15.9	12.1	11.8	10.3	13.9	11.0	8.5	19.5	18.7	•	18.2	14.4	16.5	22.7
Composition (percent by volume)	Amount of	56.3	47.2	57.5	50.0	57.9	38.5	52.5	9.75	54.2	75.9	58.7	27.7	62.2	66.1	62.9	57.7	
		Types*			M+F+y	M+F+y	M+F+y	M+F+y	M+F+y	M+F+y	М+F+ү	M+F+γ	M+F+y	M+F	Μ+F+γ	M+F+y	M+F+γ	M+F+y
ing	Tempering temperature (°C)		570	570	929	570	929	570	570	570	570	570	920	570	570	570	920	570
Quenching-tempering	hing	Cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling	Air cooling
Que	Quenching	Heating temperature (°C)	920	920	920	920	920	920	920	920	920	920	920	920	920	920	920	920
······································	Steel No.		1A	18	10	10	1E	1F	16	1H	1)	¥	11	1M	1N	1.	ō,	1R
	Steel	o Z	3-1	3-2	3-3	3-4	3-5	3-6	3-7	3-8	3-9	3-10	3-11	3-12	3-13	3-14	3-15	3-16

M: Martensite, F: Ferrite, γ : Retained austenite

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[0093] According to the examples of the present invention, the generation of cracks in the surface of the steel pipe was not observed, the yield strength YS was high, such as 654 MPa or more, the corrosion rate was also low, and no pitting occurred; hence. Hence, a steel pipe was obtained having superior hot workability and corrosion resistance in a severe corrosive environment in which CO₂ was present and the temperature was high, such as 230°C. Furthermore, since 5% or more of a ferrite phase was contained, a steel pipe was obtained having superior corrosion resistance in a severe corrosive environment in which CO₂ was present and the temperature was high, such as 230°C; a high strength, such as a yield strength of 654 MPa or more; and a high toughness having an absorption energy of 50 J or more at -40°C. In addition, as for steel pipes Nos. 13 and 14, the content of Al was high, the toughness was slightly decreased, and pitting occurred; however. However, the degree thereof was not significant, and the diameter of the pitting hole by pitting was less than 0.2 mm.

[0094] On the other hand, according to the comparative examples—which were outside the range of the present invention, cracks were generated in the surface since the hot workability was degraded; or the corrosion rate was high and pitting occurred since the corrosion resistance was degraded. In particular, in the comparative example in which the equation (2) was not satisfied, the hot workability was degraded, and as a result, scars were generated on the surface of the steel pipe. In addition, when the amount of ferrite was out of the preferable range—of the present invention, the strength was decreased, and a high strength having a yield strength of 654 MPa or more could not be achieved.

Industrial Applicability

[0095] According to the present invention, aA stainless steel pipe for use in oil wells can be stably manufactured at an inexpensive cost, the stainless steel pipe having a high strength and sufficient corrosion resistance in a severe corrosive environment in which CO₂ and Cl⁻ are present and the temperature is high, or further having a high toughness; hence. Hence, from the present invention, significant industrial advantages can be obtained. In addition, according to the present invention, another advantage can also be obtained in that a sufficient strength as an oil-well pipe can be obtained only by performing heat treatment after pipe makingpipe making.